

# (11) EP 1 506 854 A1

(12)

# **EUROPEAN PATENT APPLICATION**

(43) Date of publication: 16.02.2005 Bulletin 2005/07 (51) let CL7: **B41C 1/10**, B41N 3/00

(21) Application number: 03102530.7

(22) Date of filing, 13.08,2003

(84) Designated Contracting States:

AT BE BG CH CY CZ DE DK EE ES FI FR GB GR HU IE IT LI LU MC NL PT RO SE SI SK TR Designated Extension States:

AL LT LV MK

(71) Applicant: AGFA-GEVAERT 2640 Mortsel (BE) (72) Inventors:

- Verschueren, Eric 2640, Mortsel (BE)
- Van Damme, Marc 2640, Mortsel (BE)

# (54) Method for postbaking a lithographic printing plate

(57) A method for making a lithographic printing plate is disclosed wherein the method comprises the steps of: (i) providing a lithographic printing plate precursor comprising a photosensitive coaling on a substrate having a hydrophilic surface, (ii) image-wise exposing said photosensitive coaling, (iii) developing said exposed coaling to form an image on said substrate and optionally gumming the plate and (iv) baking the image

on the plate, wherein said baking step is carried out within a dwell-time of less than 1 minute and wherein the chemical resistance of the coating against printing liquids and press chemicals is improved. This baking step is preferably carried out by exposing the printing plate to an infrared radiation source, preferably in a dynamic configuration.

# Description

### FIELD OF THE INVENTION

5 [0001] The present invention relates to a method for preparing a lithographic printing plate.

#### BACKGROUND OF THE INVENTION

[0002] Ulthographic printing presses use a so-called printing master such as a printing plate which is mounted on a cylinder of the printing press. The master carries a lithographic image on its surface and a print is obtained by applying ink to said image and then transferring the link from the master onto a receiver material, which is typically paper, in conventional, so-called "wet" lithographic printing, ink as well as an aqueous fountain solution (also called dampening liquid) are supplied to the fithographic image which consists of oleophilic (or hydrophobic, i.e. ink-accepting, water-repetiting) areas as well as hydrophilic (or oleophobic, i.e. water-accepting, ink-repetiting) areas, in so-called driographic printing, the lithographic image consists of ink-accepting and ink-abhesive (ink-repetiting) areas and during driographic printing, only ink is supplied to the master.

Printing masters are generally obtained by the so-called computer-to-film method wherein various pre-press steps such as typetace selection, scanning, color separation, screening, trapping, tayout and imposition are accomplished digitally and each color selection is transferred to graphic arts film using an image-setter. After processing, the film can be used as a mask for the exposure of an imaging material called plate precursor and after plate processing, a printing plate is obtained which can be used as a master.

A typical printing plate precursor for computer-to-film methods comprise a hydrophilic support and an image-recording layer of a photosensitive polymer layers which include UV-sensitive diazo compounds, dichromate-sensitized hydrophilic colloids and a large variety of synthetic photopolymers. Particularly diazo-sensitized systems are widely used. Upon image-wise exposure, typically by means of a film mask in a UV contact frame, the exposed image areas become insoluble and the unexposed areas remain soluble in an equeous alkaline developer. The plate is then processed with the developer to remove the diazonium salt or diazo resin in the unexposed areas. So the exposed areas define the image areas (printing areas) of the printing master, and such printing plate precursors are therefore called 'negative-working'. Also positive-working materials, wherein the exposed areas define the non-printing areas, are known, e.g. plates having a novolac/naphtoquinone-diazide coating which dissolves in the developer only at exposed areas.

In addition to the above photosensitive meterials, also heat- or infrared-sensitive printing plate precursors have become very popular. Such materials offer the advantage of daylight-stability and are especially used in the so-called computer-to-plate method wherein the plate precursor is directly exposed. i.e. without the use of a film mask. The meterial is exposed to heat or to infrared light and the generated heat triggers a (physico-)chemical process, such as abtation, polymerization, insolubilisation by cross-linking of a polymer, heat-induced solubilisation, decomposition, or particle coagulation of a thermoplastic polymer latex.

The known heat- or infrared-consitive printing plate precursors typically comprise a hydrophilic support and a coating containing an eleophilic polymer, which is alkali-soluble in exposed areas (positive working material) or in non-exposed areas (negative working material) and an IR-absorbing compound. Such an eleophilic polymer is typically a phenolic testin.

The ink and fountain solution which are supplied to the plate during the printing process, may affack the coating and, consequently, the resistance of the coating against these liquids, hereinafter referred to as "chemical resistance", may affect the printing run length. The printing run length can be improved by heating the image formed on the support after developing and gumming process. The known process for post-baking a printing plate is carried out by heating the plate in a heating oven at a temperature of about 235°C to 270°C during a relatively long time of about 2 to 5 minutes, even upto 10 minutes. This time consuming baking process is unfavourable for those applications where a quick availability of lithographic printing plates with a high printing run length is required, e.g. in the preparation of newspaper printing plates. In this sector, the systems which are used for image-wise exposing printing plate precursors and for processing and gumming these precursors, are very complicated machines which are able to deliver upto 200 plates per hour. However, this high throughput is reduced due to the slow baking process. The speed of the baking step is substantially increased by the method as disclosed in the present invention and this method allows the preparation of lithographic printing plates with a high chemical resistance of the coating in a short period of time.

## SUMMARY OF THE INVENTION

55

[0003] It is an aspect of the present invention to provide a process for making a lithographic printing plate comprising a baking step which is carried out within a dwell-time of less than 1 minute and wherein the chemical resistance of the

coating against printing liquids and press chemicals is improved. This object is realized by the method as defined in claim 1

Specific embodiments of the invention are defined in the dependent claims

#### 5 DETAILED DESCRIPTION OF THE INVENTION

10

30

88

55

[0004] In order to obtain a linegraphic printing plate with an improved printing run length, it is important to increase the chemical resistance of the photosensitive coating against the printing liquids such as the dampening liquid and ink, and against the press chemicals such as cleaning liquids for the plate, for the blanket and for the press rollers. These printing properties are affected by a heat treatment of the printing plate after developing the image-wise exposed printing plate precursor and optionally gumming the plate. This heat treatment, hereinafter also referred to as a "baking step" or "baking process" or "baking method" or "baking", is one of the most important ways to obtain these properties in a short period of time time.

In accordance with the present invention, there is provided a method for making a lithographic printing plate comprising the steps of: (i) providing a lithographic printing plate precursor comprising a photosensitive coating on a substrate having a hydrophilic surface, (ii) image-wise exposing said photosensitive coating, (iii) developing said exposed coating to form an image on said substrate and optionally gurmming the plate and (iv) baking the image on the plate, wherein said baking step is carried out within a dwelf-time of less than 1 minute, preferably less than 30 seconds, more preferably less than 10 seconds.

It is also an aspect of the present invention that this baking step is carried out by exposing the printing plate to an infrared radiation source, hereinatter also referred to as an "infrared emitter" or shortly "emitter". An infrared radiation source may be any device or system emitting infrared radiation. The wavelength of the emitter may range between 760 nm and 6 µm, preferably between 770 nm and 5 µm, more preferably between 770 nm and 3.5 µm and most preferably between 770 nm and 2 µm. The emitter radiates substantially infrared light, without excluding side radiation of wavelengths outside the infrared wavelength range. The peak wavelength of the emitter may be in the near infrared range (i.e. between 750 nm and 1.5 µm), in the middle infrared range (i.e. between 1.5 µm and 6 µm), in the short wave infrared range (i.e. between 750 nm and 2 µm) or in the medium wave infrared range (i.e. between 2 µm and 3.5 µm); in the near or short wave infrared ranges is preferred. The emitter may be an infrared lamp. The temp may have a reflector. The reflector has preferably a parabolic or an elliptic configuration; an elliptic configuration is more preferred. Infrared emitters, suitable for the present invention, may be selected from the following emitter types:

- MICOR-KIR-BANDSTRAHLER TYP LE/LP infrared smitters, available from MICOR GmbH, Kröfteler Strasse 11, Idatein-Heftrich, Germany.
- SHORT WAVE intrared emitter, available from HERAEUS NOBLELIGHT GmbH, Reinhard-Heraeus-Ring 7, Kleinoatheim, Germany.
- INFRALIGHT-HALOGEN infrared emitter, available from HERAEUS NOSLELIGHT GmbH, Reinhard-Heraeus-Ring 7, Kleinostheim, Germany.
- CARBON infrared emiliter, available from HERAEUS NOBLELIGHT GmbH, Reinhard-Heraeus-Ring 7, Kleinostheim, Germany,
- CARBON HIGH POWER infrared emitter, available from HERAEUS NOBLELIGHT GmbH. Reinhard-Heraeus-Bing 7, Kleinostheim, Germany.
  - CARBON TWIN infrared emitter, evallable from HERAEUS NOBLELIGHT GrobH, Reinhard-Heraeus-Ring 7, Kleinostheim, Germany.

(9006) Typical initiated emitters are preferably characterised by one or more of the following specifications or configurations; the length of the emitter may vary between 150 mm and 5 m; the voltage of the emitter may vary between 57.5 V and 600 V, the power output at maximum current may vary between 180 W and 15 kW; the mean linear power density may vary between 16 W/cm and 70 W/cm; the maximum power density may vary between 20 kW/m² and 190 kW/m²; the filament temperature of the emitter may up to 2200°C; the emitter may have a parabolic or elliptic reflector, optionally with a gold coating; the emitter may have a one-side or two-side connection; the emitter may have a twin tube emitter; the working of the emitter may be controlled, timed and/or adjusted by means of a control system.
[9006] In accordance with a specific embodiment of the present invention, the baking step is cerried out by exposing

tweet in accordance with a specific embodiment of the present invention, the baking step is carried but by exposing the printing plate to at least one of said infrared emitting lamps. This baking process may be carried out in a static or dynamic configuration, a dynamic configuration is preferred.

In accordance with a specific embodiment of the present invention, the beking step is carried out by the following steps: radiating the printing plate to an emitter, and transporting the printing plate relative to the emitter. These two steps may be carried out separately, for example, transporting the plate relative to the emitter and then radiate the plate, or, by preference, these two steps are carried out simultaneously, for example, rediating the plate while trans-

porting the plate relative to the emitter. In these configurations, at least one emitter is used, but two or more emitters of the same or different type, optionally in a serial configuration, may also used. The specifications and configuration of each emitter are chosen corresponding with the configuration wherein the baking step is carried out, such that the plate is homogeneously heated and sufficiently high in order to obtain a high chemical resistance.

9

18

40

In a dynamic configuration, the effective time of radiation of the plate and the intensity of radiation are important parameters for the efficiency of the baking process. The effective time of radiation of the plate may be defined for each emitter by the dwell-time. The dwell-time is the ratio of the exposure breadness of the emitter over the relative transporting speed of the plate to the emitter. The exposure breadness of the emitter may depend on the type of the emitter, the type of the reflector and the distance to the plate. Depending on the intensity of radiation, the relative transporting speed of the plate to the emitter(s) may vary between 0.5 m/minute and 20 m/minute; more specific between 1 m/minute and 15 m/minute, most specific between 2 m/minute and 10 m/minute. The value of the dwell-time for each emitter is lower than 1 minute, preferably lower than 30 seconds, more preferably lower than 10 seconds.

In this dynamic configuration the printing plate may be mounted in an isolated manner to prevent heat conduction to the environment. The printing plate may preferably be isolated by mounting a Tellon-material, such as a foll or a web, between the plate and the transporting means.

In order to obtain a high chemical resistance, the coating on the plate exceeds a threshold temperature during the heating step. The threshold temperature depends on the type of the coating, more specific on the type of the chemical compounds used in the coating, and may also depend on the type of the surface of the support of the plate or on the type of a subbing layer present between the support and the coating. A typical example of such a threshold temperature may be a value above 100°C or above 150°C or even more above 200°C. The temperature of the coating may not exceed an upperlimit value where the coating starts decomposing, degrading or carbonating or where the coating starts ablating of the support. A typical example of an upperlimit temperature may be about 800°C, more precisely about 600°C. Typically, but not limited thereto, the coating of the plate may reach a temperature of above 120°C, more specific above 150°C, even more specific above 200°C. In order not to exceed the upperlimit temperature of the coating it is important to chose, in correspondence with the selected dwell-time for each emitter, the specifications and configuration of the emitter(s).

[0007] In another preferred embodiment of the present invention, the baking process comprises the following steps radiating the printing plate while the printing plate is transported to the emitter wherein the emitter is mounted in a fixed manner and the plate transports during the radiation. The plate may be transported in any direction, preferably in a horizontal direction, a vertical direction or a tilt direction; a horizontal direction is preferred. The position of the emitter may be preferably mounted above or beside the plate depending on the chosen transport direction (horizontally, vertically or tilt). The most preferred configuration is a horizontal transport of the plate with an emitter positionned above the coated side of the plate.

By the use of an emitter with an elliptic reflector the highest intensity of radiation may be obtained at the focus distance of the elliptic reflector. The printing plate may be very efficient heated when the coating of the plate is placed at this focus distance to the emitter; this configuration is preferred when an elliptic reflector is used in the baking step.

In order to obtain a shorter period of time for parrying out the baking step, resulting in a high chemical resistance of the coating, the specifications and configurations for each emitter are selected in such a way that the intensity of radiation on the coating in correspondence with the relative speed of plate to the emitter is sufficiently high such that the coating reaches its threshold temperature, in accordance to the present invention, the baking step is carried out within a dwell-time of less than 1 minute, preferably less than 30 seconds and more preferably less than 10 seconds.

In another embodiment of the present invention, an apparatus for baking an image, obtained after developing an image-wise exposed photosensitive coating on a substrate having a hydrophilic surface, is disclosed, wherein said apparatus comprises an infrared radiation source, characterised in that said infrared radiation source is a near infrared temp or a middle infrared lamp. Configurations as described above for dynamic and static configurations may preferably used in this apparatus. In accordance to another embodiment of the present invention, a processor for developing an image-wise exposed photosensitive coating on a substrate having a hydrophilic surface, is disclosed, wherein the processor comprises the apparatus for baking an image as disclosed above. The baking apparatus may be preferably integrated in a processor. In this configuration, the exposed photosensitive coating is developed and, after developing is further transported relative to the infrared radiation source. The baking process is carried out in a static or dynamic configuration; the dynamic configuration is preferred. The baking process is carried out within a dwell-time of less than 1 minute. In a dynamic configuration, the baking process is carried out within a dwell-time varying from 1 to 50 seconds depending on the processor speed and the type of emitter(s) which is (are) used.

The baking process may be also carried out in a static configuration by the use of one or more emitters. In a preferred configuration the emitter or emitters are positionned above the coating of the plate. The specifications and configuration of each emitter are chosen in corresponding with the intensity of radiation, needed to heat the plate homogeneously and sufficiently high in order to obtain a high chemical resistance of the coating. In the static configuration the effective time of radiation of the plate is defined as the dwell-time.

[0008] In another embodiment of the present invention, the baking step is carried out by exposing the printing plate to an infrared radiation source in combination with an ultraviolet radiation source. The ultraviolet radiation source is preferably an UV lamp. The infrared lamp in this baking process may be combined with an UV lamp in order to further improve the efficiency of the baking process. Under this combination any configuration is understood whereby the printing plate is simultaneously or successively exposed by an IR radiation source and an UV radiation source.

Types of UV radiation sources which can be used in these combinations are Medium Pressure Mercury Arc lamps, with a power range of 1kW to 31kW, Metal Halide lamps, doped with Gallium, Metal Halide lamps, doped with iron, or Excimer UV lamps. Examples of such UV radiation sources are the AMBA UV Curing Lamps, commercially available from HERAEUS NOBLELIGHT GmbH, Reinhard-Heraeus-Ring 7, Kleinoetheim, Germany.

[0009] This chemical resistance of the coating can be measured by the test as described in the examples.

5

[0010] The lithographic printing plate precursor may comprise a photosensitive coating on a substrate having a hydrophilic surface. The photosensitive coating may comprise at least one image-sensitive layer which may be sensitive to UV light, violet light, visible light or infrared light. The image-sensitive layer of the photosensitive coating may comprise at least a light absorbing compound corresponding with the wavelength of the light used by the image-wise recording. Optionally, in addition to this light absorbing compound, the photosensitive coating may further comprise an infrared absorbing compound, suitable for absorbing IR-radiation during the baking step. This infrared absorbing compound may increase the efficiency of the baking process.

The photosensitive coaling may also be sensitive to heat. Typical for such a heat-sensitive coaling is that the image-sensitive layer comprises an intrared light absorbing compound, which, on image-wise exposing to infrared light, is able to convert the absorbed infrared light into heat. According to a preferred embodiment, the photosensitive coating is sensitive to infrared light or to heat and this coating comprises preferably an infrared absorbing compound.

The photosensitive coating provided on the hydrophilic support, is preferably sensitive to infrared light or heat and can preferably be handled in normal working lighting conditions (daylight, fluorescent light) for several hours. This coating preferably does not contain UV-sensitive compounds which have an absorption maximum in the wavelength range of 200 nm to 400 nm such as diazo compounds photoscida, photoinitiators, quinone diazides, or sensitizers. Preferably the coating neither contains compounds which have an absorption maximum in the blue and green visible light wavelength range between 400 and 600 nm.

[0011] The photosonalitive coating may be composed of at least one image-sensitive layer, hereinafter also referred to as image-recording layer. The coaling may also contain one or more additional layer(s), adjacent to the imagerecording layer. Besides the layers discussed hereafter, the coating may further comprise e.g., a "subbing" layer which improves the adhesion of the coating to the support, a covering layer which protects the coating against contamination or mechanical damage, and/or a light-to-heat conversion tayer which comprises an infrared light absorbing compound, [0012] After exposure, the material is developed. "Developing" and "processing" are used herein as equivalent terms, Development can be carried out by supplying to the coating a liquid comprising a hydrophilic phase, thereby removing the coating from the support at exposed or non-exposed areas, depending on the type of the plate, namely positive or negative working. Said liquid can be selected from the group consisting of water, an aqueous liquid, gum, fountain and single-fluid ink. The meterial can be developed by supplying fountain and/or printing ink, preferably by supplying first fountain and subsequently ink. This method is preferably used in combination with an on-press exposure step. Another development method, also suitable for on-press development, especially in driographic presses, is performed by supplying single-fluid ink. Single-fluid inks which are suitable for use in the method of the present invention have been described in US 4,045,232 and US 4,981,517. A suitable single-fluid ink comprises an ink phase, also called the hydraphotic or eleophilic phase, and a polyol phase as described in WO 00/32705. More information on the development with single-fluid ink can be found in EP-A no. 01000633, filed on 15.11.2001.

When exposed in an oil-press plate-setter, the material can be processed on-press by supplying ink and/or lountain as mentioned before or off-press, e.g. by supplying water, an equenus liquid or a gum solution.

After the development or simultaneously with the development, the plate is optionally gummed. A gum solution is typically an aqueous liquid which comprises one or more surface protective compounds that are capable of protecting the lithographic image of a printing plate against contamination or damaging. Suitable examples of such compounds are film-forming hydrophilic polymers or surfactants. More information on the development with a gum solution can be found in EP-A no. 02100228, filed on 06.03.2002.

After developing or gurmling, the plate can be dried before the beking process or can be dried during the baking process likelii.

[0013] In an example of a suitable negative-working alkaline developing printing plate, the coating comprises a phenotic resin and a latent Bränsted acid which produces acid upon heating or IR radiation. These acids catalyze crosslinking of the coating in a post-exposure heating step and thus hardening of the exposed regions. Accordingly, the non-exposed regions can be washed away by a developer to reveal the hydrophilic substrate underneath. For a more detailed description of such a negative-working printing plate precursor we refer to US 6,255,042 and US 6,063,544 and to references cited in these documents.

[0014] In an example of a positive-working lithographic printing plate precursor, the coating is capable of heat-induced solubilization, i.e. the coating is resistant to the developer and ink-accepting in the non-exposed state and becomes soluble in the developer upon image-wise exposure to such an extent that the hydrophilic surface of the support is revealed thereby.

The coating may contain at least one polymeric binder that is soluble in an aqueous alkaline developer. Preferred polymers are phenolic resins, e.g. nevolae, resoles, polyvinyl phenois and carboxy-substituted polymers. Typical examples of such polymers are described in DE-A-4007428, DE-A-4027301 and DE-A-4446820.

5

30

25

In a preferred positive-working lithographic printing plats precursor, the coating also contains one or more dissolution inhibitors. Dissolution inhibitors are compounds which reduce the dissolution rate of the hydrophobic polymer in the aqueous alkaline developer at the non-exposed areas of the coating and wherein this reduction of the dissolution rate is destroyed by the heat generated during the exposure so that the coating readily dissolves in the developer at exposed areas. The dissolution inhibitor exhibits a substantial latitude in dissolution rate between the exposed and non-exposed areas. By preference, the dissolution inhibitor has a good dissolution rate latitude when the exposed coating areas have dissolved completely in the developer before the non-exposed areas are attacked by the developer to such an extent that the ink-accepting capability of the coating is affected. The dissolution inhibitor(s) can be added to the layer which comprises the hydrophobic polymer discussed above.

The dissolution rate of the non-exposed coating in the developer is preferably reduced by interaction between the hydrophobic polymer and the inhibitor, due to e.g. hydrogen bonding between these compounds. Suitable dissolution inhibitors are preferably organic compounds which comprise at least one aromatic group and a hydrogen bonding site, e.g. a carbonyl group, a sulfonyl group, or a nitrogen atom which may be queternized and which may be part of a heterocyclic ring or which may be part of an amino substituent of said organic compound. Suitable dissolution inhibitors of this type have been disclosed in e.g. EP-A 825927 and 823327.

Water-regellent polymers represent an another type of suitable dissolution inhibitors. Such polymers seem to increase the developer resistance of the coating by repelling the aqueous developer from the coating. The waterrepellent polymers can be added to the tayer comprising the hydrophobic polymer and/or can be present in a separate layer provided on top of the layer with the hydrophobic polymer, in the latter embodiment, the water-repellent polymer forms a barrier layer which shields the coating from the developer and the solubillly of the barrier layer in the developer or the penetrability of the barrier layer by the developer can be increased by exposure to heat or infrared light, as described in e.g. EP-A 864420, EP-A 950517 and WO99/21725. Preferred examples of the water-repellent polymers are polymers comprising siloxane and/or perfluoroalkyl units. In one embodiment, the coating contains such a waterrepellent polymer in an amount between 0.5 and 25 mg/m², preferably between 0.5 and 15 mg/m² and most preferably between 0.5 and 10 mg/m². When the water-repellent polymer is also ink-repelling, s.g. in the case of polysiloxanes, higher amounts than 25 mg/m² can result in poor ink-acceptance of the non-exposed areas. An amount lower than 0.6 mg/m² on the other hand may lead to an unsatisfactory development resistance. The polysiloxane may be a linear, cyclic or complex cross-linked polymer or copolymer. The term polysiloxane compound shall include any compound which contains more than one siloxane group -Si(R,R')-O-, wherein R and R' are optionally substituted alkyl or aryl groups. Preferred silications are phenylalkylalloxanes and dialkylalloxanes. The number of silicatine groups in the (co) polymer is at least 2, preferably at least 10, more preferably at least 20, it may be less than 100, preferably less than 60. In another embodiment, the water-repellent polymer is a block-copolymer or a graft-copolymer of a poly(alkylene oxide) block and a block of a polymer comprising sitoxane and/or perfluoroaliky) units. A sulfable copolymer comprises about 15 to 25 siloxane units and 50 to 70 alkylone oxide groups. Preferred examples include copolymers comprising phenylmethylelloxane and/or d/methylsiloxane as well as ethylene oxide and/or propytene oxide, such as Tego Glide 410, Tego Wet 265, Tego Protect 5001 or Silikophen P50/X, all commercially available from Tego Chemie, Essen, Germany. Such a copolymer acts as a surfactant which upon coating, due to its bitunctional structure, automatically positions itself at the interface between the coating and air and thereby forms a separate top layer even when the whole coating is applied from a single coating solution. Simultaneously, such surfactants act as a spreading agent which improves the coating quality. Alternatively, the water-repellent polymer can be applied in a second solution, coated on top of the layer comprising the hydrophobic polymer. In that embodiment, it may be advantageous to use a solvent in the second coating solution that is not capable of dissolving the ingredients present in the first laver so that a highly concentrated water-repellent phase is obtained at the top of the coating.

Preferably, also one or more development accelerators are included in the coeting, i.e. compounds which act as dissolution promoters because they are capable of increasing the dissolution rate of the non-exposed coeting in the developer. The simultaneous application of dissolution inhibitors and accelerators allows a precise fine funing of the dissolution behavior of the coating. Suitable dissolution accelerators are cyclic acid anhydrides, phenols or organic acids. Examples of the cyclic acid anhydride include phthalic anhydride, tetrahydrophthalic anhydride, maleic anhydride, tetrachlorophthalic anhydride, maleic anhydride, chloromateic anhydride, alpha -phenylmaleic anhydride, succinic anhydride, and pyromellitic anhydride, as described in U.S. Patent No. 4,115,128. Examples of the phenols include bisphenol A, p-nitrophenol, p-ethoxyphenol, 2,4,4°-trihydroxybenzophenone, 2,3,4-trihydroxy-benzophenone.

phenone, 4-hydroxybenzophenone, 4,4',4"-trihydroxy-triphenylmethene, and 4,4',3",4"-tetrahydroxy-3,5,3',5'-tetramethyltriphenyl-methane, and the like. Examples of the organic acids include sulfonic acids, sulfinic acids, alkylsulfuric acids, phosphates, and carboxylic acids, as described in, for example, JP-A Nos. 60-88,942 and 2-96,766. Specific examples of these organic acids include p-toluenesulfonic acid, dedecylbenzenesulfonic acid, p-toluenesulfinic acid, athylsulfuric acid, phenylphosphanic acid, phenylphosphate, benzoic acid, isophthelic acid, adipic acid, p-toluic acid, 3,4-dimethoxybenzoic acid, phthalic acid, terephthelic acid, 4-cyclohexene-1,2-dicarboxylic acid, erucic acid, lauric acid, n-undecanoic acid, and ascorbic acid. The amount of the cyclic acid anhydride, phenol, or organic acid contained in the coating is preferably in the range of 0.05 to 20% by weight, relative to the coating as a whole.

[0015] In another preferred positive-working lithographic printing plate precursor, the coating further contains hydrophobic thermoplastic polymer particles. Specific examples of suitable hydrophobic polymers are e.g. polyethylene, poly(vinyl chioride), poly(meth)acrylete), poly(ethyl (meth)acrylete), poly(vinylidene chloride), poly(meth)acrylete), poly(ethyl (meth)acrylete), poly(vinylidene chloride), poly(meth)acrylete), poly(meth)acrylete), poly(ethyl (meth)acrylete), poly(vinylidene chloride), poly(meth)acrylete), poly(meth)acrylete), poly(ethyl (meth)acrylete), poly(ethyl (meth)acrylete), poly(ethyl (meth)acrylete), poly(ethyl (meth)acrylete), poly(meth)acrylete), poly(ethyl (meth)acrylete), poly(ethyl (meth)acrylete), poly(ethyl (meth)acrylete), poly(meth)acrylete), poly(ethyl (meth)acrylete), poly(ethyl (meth)acrylete), poly(ethyl (meth)acrylete), poly(meth)acrylete), poly(ethyl (meth)acrylete), poly(meth)acrylete), poly(ethyl (meth)acrylete), poly(ethyl (meth)acry

The weight average molecular weight of the thermoplastic polymer particles may range from 5,000 to 1,000,000 g/mol. The hydrophobic particles preferably have a number average particle diameter below 200 nm, more preferably between 10 and 100 nm. The amount of hydrophobic thermoplastic polymer particles contained in the image-recording layer is preferably between 20 wt.% and 55 wt.% and more preferably between 25 wt.% and 55 wt.% and more preferably between 30 wt.% and 45 wt.%.

The hydrophobic thermoplastic polymer particles can be provided as a dispersion in an aqueous coating liquid of the image-recording layer and may be prepared by the methods disclosed in US 3,476,937. Another method especially suitable for preparing an aqueous dispersion of the thermoplastic polymer particles comprises:

- dissolving the hydrophobic thermoplastic polymer in an organic water immiscible solvent,
- dispersing the thus obtained solution in water or in an aqueous medium and
- removing the organic solvent by evaporation.

5

25

30

39

40

48

50

The image-recording layer further may comprise a hydrophilic binder, e.g. homopolymers and copolymers of vinyl alcohol, acrylamide, methylol acrylamide, methylol methacrylamide, acrylic acid, hydroxyethyl acrylate, hydroxyethyl methacrylate or maleic anhydride/vinylmethylether copolymers. The hydrophilicity of the (co)polymer or (co)polymer mixture used is preferably the same as or higher than the hydrophilicity of polyvinyl acetate hydrolyzed to at least an extent of 60 percent by weight, preferably 80 percent by weight. Sinders with carboxylic pendant groups, e.g. poly(meth)acrylic acid, are preferred.

Due to the heat generated during the exposure step, the hydrophobic thermoplastic polymer particles fuse or coagulate so as to form a hydrophobic phase which corresponds to the printing areas of the printing plate. Coagulation may result from heat-induced coalescence, softening or melting of the thermoplastic polymer particles. There is no specific upper limit to the coagulation temperature of the thermoplastic hydrophobic polymer particles, however the temperature should be sufficiently below the decomposition temperature of the polymer particles. Preferably the coagulation temperature is at least 10°C below the temperature at which the decomposition of the polymer particles occurs. The coagulation temperature is preferably higher than 50°C, more preferably above 100°C.

After developing and optionally gumming, the plate can be dried and baked. The baking process can proceed at a temperature above the coagulation temperature of the thermoplastic polymer.

[0016] The image-recording layer may also contain other ingredients such as additional binders, surfactants, colorants, development inhibitors or accelerators, and especially one or more compounds that are capable of converting infrared light into heat.

[0017] According to a more preferred embodiment, the lithographic printing plate precursor is image-wise exposed to infrared light, which is converted into heat by an infrared light absorbing agent, which may be a dye or pigment having an absorption maximum in the infrared wavelength range. The concentration of the sensitizing dye or pigment in the coating is typically between 0.25 and 10.0 wt.%, more preferably between 0.5 and 7.5 wt.% relative to the coating as

a whole. Preferred IR-absorbing compounds are dyes such as cyanine or mercoyanine dyes or pigments such as carbon black. A suitable compound is the following intrared dye:

[0018] The coating may further contain an organic dye which absorbs visible light so that a perceptible image is obtained upon imagewise exposure and subsequent development. Such a dye is often called contrast dye or indicator dye. Preferably, the dye has a blue color and an absorption maximum in the wavelength range between 600nm and 750 nm. Although the dye absorbs visible light, it preferably does not sensitize the printing plate precursor, i.e. the coating does not become more soluble in the developer upon exposure to visible light, Suitable examples of such a contrast dye are the quaternized triarylmethane dyes. Another suitable compound is the following dye:

25

50

55

[0019] The infrared light absorbing compound and the contrast dye may be present in the layer comprising the hydrophobic polymer, and/or in the barrier layer discussed above and/or in an optional other layer. According to a highly preferred embodiment, the infrared light absorbing compound is concentrated in or near the barrier layer, e.g. in an intermediate layer between the layer comprising the hydrophobic polymer and the barrier layer.

[0020] The support has a hydrophilic surface or is provided with a hydrophilic layer. The support may be a sheetlike material such as a plate or it may be a cylindrical element such as a sleeve which can be slid around a print cylinder of a printing press. Preferably, the support is a metal support such as aluminum or stainless steet.

[0021] A particularly preferred lithographic support is an electrochemically grained and anodized aluminum support. Graining and anodizing of aluminum lithographic supports is well-known. The grained aluminum support used in the material of the present invention is preferably an electrochemically grained support. The acid used for graining can be e.g. nitric acid. The acid used for graining preferably comprises hydrogen chloride. Also mixtures of e.g. hydrogen chloride and ecetic sold can be used.

The grained and enodized aluminum support may be post-treated to improve the hydrophilic properties of its surface. For example, the aluminum support may be silicated by treating its surface with a sodium silicate solution at elevated temperature, e.g. 95°C. Alternatively, a phosphate treatment may be applied which involves treating the alu-

minum exide surface with a phosphate solution that may further contain an inergenic fluoride. Further, the aluminum exide surface may be rinsed with an organic acid and/or salt thereof, e.g. carboxylic acids, hydroxycarboxylic acids, sulfenic acids or phosphenic acids, or their salts, e.g. succinates, phosphates, phosphenes, sulfates, and sulfenites. A citric acid or citrate solution is preferred. This treatment may be carried out at room temperature or may be carried out at a slightly devated temperature of about 30 to 50°C. A further post-treatment involves thising the aluminum exide surface with a bicarbonate solution. Still further, the aluminum exide surface may be treated with polyvinylphosphenic acid, polyvinylphosphenic acid, polyvinyl alcohol, polyvinylsulfenic acid, polyvinylphosphenic acid, polyvinyl alcohol, and acetals of polyvinyl alcohols formed by reaction with a sulfenited alignatic alidehyde. It is further evident that one or more of these post-treatments may be carried out alone or in combination. More detailed descriptions of these treatments are given in GB-A- 1 084 070, DE-A- 4 423 140, DE-A- 4 417 907, EP-A- 859 909, EP-A- 537 633, DE-A- 4 001 466, EP-A- 292 801, EP-A- 291 750 and US-P- 4 488 005. [0022] According to another embodiment, the support can also be a flexible support, which is provided with a hydrophilic layer, hereinafter called base layer. The flexible support is e.g. paper, plastic film, thin aluminum or a laminate thereof. Preferred examples of plastic film are polyethylene terephthalate film, polyethylene naphthalate film, callulose acetate film, polyetyrene film, polyethylene film, callulose acetate film, polyetyrene film, polyethylene film, callulose

30

18

25

48

The base layer is preferably a cross-linked hydrophilic layer obtained from a hydrophilic binder cross-linked with a hardening agent such as formaldehyde, glyoxal, polyisocyanate or a hydrofyzed letra-alkylorthosilicate. The latter is particularly preferred. The thickness of the hydrophilic base layer may very in the range of 0,2 to 25 µm and is preferably 1 to 10 µm.

The hydrophilic binder for use in the base layer is e.g. a hydrophilic (co)polymer such as homopolymers and copolymers of vinyl alcohol, acrylamide, methylol acrylamide, methylol methacrylamide, acrylic acid, methacrylic acid, hydroxyethyl acrylate, hydroxyethyl methacrylate or maleic anhydride/vinylmethylether copolymers. The hydrophilicity of the (co)polymer or (co)polymer mixture used is preferably the same as or higher than the hydrophilicity of polyvinyl acetate hydrolyzed to at least an extent of 60% by weight, preferably 80% by weight.

The amount of hardening agent, in particular tetraalkyl orthosilicate, is preferably at least 0.2 parts per part by weight of hydrophilic binder, more preferably between 0.5 and 5 parts by weight, most preferably between 1 parts and 3 parts by weight.

The hydrophilic base layer may also contain substances that increase the mechanical strength and the porosity of the layer. For this purpose colloidal silica may be used. The colloidal silica employed may be in the form of any commercially available water dispersion of colloidal silica for example having an average particle size up to 40 nm, e.g. 20 nm. In addition inert particles of larger size than the colloidal silica may be added e.g. silica prepared according to Stöber as described in J. Colloid and Interface Sci., Vol. 26, 1968, pages 62 to 69 or alumina particles or particles having an average diameter of at least 100 nm which are particles of titanium dioxide or other heavy metal oxides. By incorporating these particles the surface of the hydrophilic base layer is given a uniform rough texture consisting of microscopic hills and valleys, which serve as storage places for water in background areas.

Particular examples of sulfable hydrophilic base layers for use in accordance with the present invention are disclosed in EP-A-601 240, GB-P-1 419 512; FR-P- 2 300 354, US-P- 3 971 560, and US-P- 4 284 705.

It is particularly preferred to use a fifth support to which an adhesion improving layer, also called support layer, has been provided. Particularly suitable adhesion improving layers for use in accordance with the present invention comprise a hydrophilic binder and colloidal allica as disclosed in EP-A- 619 524, EP-A- 620 502 and EP-A- 619 525. Preferably, the amount of silica in the adhesion improving layer is between 200 mg/m² and 750 mg/m². Further, the ratio of silica to hydrophilic binder is preferably more than 1 and the surface area of the colloidal silica is preferably at least 300 m²/gram, more preferably at least 500 m²/gram.

[0023] The printing plate precursor of the present invention can be exposed to infrared light with LEDs or a laser. Preferably, a laser emitting near infrared light having a wavelength in the range from about 750 to about 1500 nm is used, such as a semiconductor laser diode, a Nd:YAG or a Nd:YLF laser. The required laser power depends on the sensitivity of the image-recording layer, the pixel dwell time of the taser beam, which is determined by the spot diameter (typical value of modern plate-setters at 1/e<sup>2</sup> of maximum intensity: 10-25 µm), the sean speed and the resolution of the exposure apparatus (i.e. the number of addressable pixels per unif of linear distance, often expressed in dots per inch or obj; typical value: 1000-4000 dpi).

[0024] Two types of leser-exposure apparatuses are commonly used: internal (FTD) and external drum (XTD) plate-setters. ITD plate-setters for thermal plates are typically characterized by a very high scan speed up to 1500 m/sec and may require a laser power of several Watts. The Agla Galileo T is a typical example of a plate-setter using the ITD-technology. XTD plate-setters operate at a lower scan speed typically from 0.1 m/sec to 10 m/sec and have a typical leser-output-power per beam from 20 mW up to 500 mW. The Creo Trendsetter plate-setter family and the Agia Excellibur plate-setter family both make use of the XTD-technology.

The known plate-setters can be used as an off-press exposure apparatus, which offers the benefit of reduced press down-time. XTD plate-setter configurations can also be used for on-press exposure, offering the benefit of im-

mediate registration in a multi-color press. More technical details of on-press exposure apparatuses are described in e.g. US 5.174,205 and US 5,163.368.

In the development step, the non-image areas of the coating can be removed by immersion in an aqueous alkaline developer, which may be combined with mechanical rubbing, e.g. by a rotating brush. The developer preferably has a pH above 10, more preferably above 12. The development step may be followed by a rinsing step, a gumming step, a drying step and/or a post-baking step.

[0025] The printing plate thus obtained can be used for conventional, so-called wet offset printing, in which ink and an aqueous dampening liquid is supplied to the plate. Another suitable printing method uses so-called single-fluid ink without a dampening liquid. Single-fluid ink consists of an ink phase, also called the hydrophobic or eleophilic phase, and a polar phase which replaces the aqueous dampening liquid that is used in conventional wot offset printing. Suitable examples of single-fluid inks have been described in US 4,045,232; US 4,981,517 and US 6,140,392. In a most preferred embodiment, the single-fluid ink comprises an ink phase and a polyol phase as described in WO 00/32705.

#### EXAMPLES

5

10

18

25

30

33

40

Preparation of the lithographic printing plate precursor TG, a negative-working thermal plate, sensitised for 830 nm;

[0028] Onto an electrochemically roughoned and anodically oxidized aluminum sheet, the surface of which has been rendered hydrophilic by treatment with an aqueous solution of polyvinyl phosphonic acid (oxide weight 3 g/m²) an image-recording layer was coated from an aqueous coating solution at a wet thickness of 30 g/m². After drying, the tayer consisted of:

- 600 mg/m<sup>6</sup> of particles of a copolymer of styrene and acrylonitrile (weight ratio 60/40), having an average particle size of 65 nm and stabilized with an anionic wetting agent;
- 80 mg/m² of infrared absorbing Dye 1,
- and 120 mg/m² of polyacrylic acid (Glascot D15 from Allied Colloids; molecular weight 2.7x107 g/mole).

# [0027] The infrared absorbing Dye 1 is

SO<sub>5</sub>O SO<sub>3</sub>H

[0028] The plate precursor thus obtained was exposed with a Creo Trendsetter (plate-setter available from Creo, Burneby, Canada), operating at 330 mJ/cm² and 150 rpm. After imaging, the plate was developed with RC620 (commercially available gumming solution from Agfa).

Preparation of the lithographic printing plate precursor TS-1, a positive-working thermal plate, sensitived for 830 nm:

- [0029] A coating solution was prepared by mixing the following ingredients:
  - 209.20 g jetrabydrofurac
  - 103.25 g of a 40.4 % by weight solution of the phenoiic polymer ALNOVOL SPN452 in Doward PM, obtained from CLARIANT GribH
- 55 410,80 g Dowanoi PM
  - 266.03 g methyl ethyl ketone
  - 2.103 g of the infrared dye S0094, commercially available from Wolfen GmbH.
  - 0.53 g of a solution of Basonyl Blue 840, commercially available from BASF

- 2.13 g of a solution of Tego Glide 410 in a concentration of 10 % by weight in Dowano! PM.
- 0.85 g of a solution of Tego Wet 265 in a concentration of 10 % by weight in Dowanot PM
- 5.34 g g of 3,4.5-irimethoxyclenamic acid.
- 5 [0030] Dowanol FM is propyleneglycol-monomethylether, a trade mark product of Dow Chemical Company. The chemical structure of \$9094 is equal to IR-1. Basonyl Blue 640 is a quaternary triarylmethane dye. TegeGlide 410 is a copolymer of polysiloxane and poly(aikylene oxide), commercially available of Tego Chemie Service GmbH.
  - TegeWet 285 is a copolymer of polysiloxane and poly(alkylene oxids), commercially available of Tego Chemie Service GmbH.

[9031] The coating solution was coated on an electrochemically grained and anodized aluminum substrate at a wet thickness of 25µm and dried at a temperature of 130°C.

Ехровить:

18

10

[0092] The printing plate precureor was exposed on a CreoScilex Trendseiter 3244 at the energy density of 150 mJ/cm²

Processing:

20

[0033] The imagewise exposed printing plate precursor was processed and gummed in an Agfa Autolith'T processor, operating at a speed of 0.96 m/min and at 26°C, and using Agfa TD5000 as developer and RC510, commercially available from AGFA, as gum.

25 Preparation of the littingraphic printing plate precursor TS-2, a positive-working thermal plate, sensitised for 1060 nm.

[0034] A coating solution was prepared by mixing the following ingredients:

- 209.19 g tetrahydrofuran
- 103.32 g of a 40.4 % by weight solution of the phenolic polymer ALNOVOL SPN452 in Doward PM, obtained from CLARIANT GmbH
  - 385.60 g Dowanol PM
  - 266.24 g methyl ethyl ketone
  - 0.614 g of the infrared Dvs 2
- 1.41 g of a solution of Basonyl Blue 840, commercially available from BASF
  - 20.15 g of a solution of Tego Glide 410 in a concentration of 1 % by weight in Doward PM.
  - 7.86 g of a solution of Tago Wei 285 in a concentration of 1 % by weight in Doward PM
  - 5.61 g g of 3,4,5-trimethoxybenzoic sold.
- 49 [0035] The intrared absorbing Dye 2 is

55

48

50

[0036] The coating solution was coated on an electrochemically grained and anodized aluminum substrate at a wet thickness of 26µm and dried at a temperature of 120°C.

## Exposure:

5

10

18

20

25

89

[8037] The printing plate precursor was exposed on a Agla Gallieo T at the energy density of 50 mil/om².

# Processing:

[0038] The imagewise exposed printing plate precursor was processed and gummed in an Agfa Autolith T processor, operating at a speed of 0.96 m/min and at 25°C, and using Agfa TD5000 as developer and RC510, commercially available from AGFA, as gum.

Preparation of the lithographic printing plate precursor VPP, a negative-working violet photpolymer plate, sensitised for 390 - 420 nm.

Preparation of the photosensitive coating:

[0039] A composition was prepared (ow = parts per weight; wt.% = weight percentage) by mixing the ingredients as specified in Table 1. This composition was coated on an electrochemically roughened and anotherally oxidized aluminum sheet, the surface of which has been rendered hydrophilic by treatment with an equeous solution of polyvinyl phosphonic acid (oxide weight 3 g/m²) and was dried at 105°C. The resulting thickness of the layer is 1.5 g/m².

Table 1:

Component	Parts per weight (g
a solution containing 32.4 wt.% of a methacrylate/ methacrylic acid copolymer (ratio methylmethacrylate: methacrylic acid of 4:1 by weight; acid number: 110 mg KOH/g) in 2-butanone (viscosity 105 mm²/s at 25°C).	0.58.8
a solution containing 88.2 wt.% of a reaction product from 1 mole of 2,2,4-trimethylhexamethylenediisocyanete and 2 moles of hydroxyethylmethacrylate (viscosity 3.30 mm²/s at 25°C)	\$.440
1,4-distyryl-3,4,5-trimethoxybenzene	0.320
Heliogene blue D 7490% dispersion (9.9 wt.%, viscosity 7.0 mm²/s at 25 °C), trade name of BASE AG	7.200

Table 1: (continued)

Component	Parts per weight (g)
2,2'-bls(2-chlorophenyi)-4,4',5,5'-tetraphenyi-1,2-bisimidazota	0.320
2-mercaptobenzothiazole	0.027
Edaplan LA 411® (1 % in Dowanol PM®, tradé mark of Dow Chemical Company)	0.900
2-butanone	47.84
Propyleneglycol-monomethylether (Doward) PM®, trade mark of Dow Chemical Company)	79.33

Preparation of the protective overcoat layer:

[0040] On top of the photosonsitive layer a solution in water containing

- 4.9 wt.% of the following composition was coated and dried at 120°C;
- 1.0 pw of partially hydrolyzed polyvinytalcohol (degree of hydrolysis 86 %, viscosity 8 mPa,s in a solution of 4 wt. % at 20 °C).
- 1.0 pw of fully hydrolyzed polyvinylalcohol (degree of hydrolysis 98 %, viscosity 4 mPa.s in a solution of 4 wt.% at 20 °C)
- 0.5 pw of polyvinylpyrrolidone (k-value 30).
- 25 The top coat had a dry thickness of 2.0 g/m².

# Exposure:

5

10

18

20

89

50

[0041] The printing plate precursor was exposed on an experimental violet platesetter device (flat bed system), equipped with a violet laser diode emitting between 392 - 417 nm. The following imaging conditions were used:

Scanning speed: 1000 m/sec
 Energy density: 50 µJ/cm²

Spot diameter : 20 μm
 Addressability : 1270 dpi

### Processing:

[0042] After imaging the plate was processed in a Agra VSP85 processor at a speed of 1,2 m/min. During the processing the plate is heated is, pro-heat step (temperature at the back of the plate is 104°C), next the protective overcost is washed off with water and the photologyer is processed in a water based alkaline developer (Agra EN 231C) at 28 °C. After a water rinsing and gumming step with RC610 the printing plate is ready.

Procedure for baking process by a dynamic IR-exposing, hereinafter referred to as "Dynamic IR":

[0043] The printing plate precursors TG, TS-1, TS-2 and VFP, as prepared above, were heated with a MI-COR-KIR-KASSETTE 36 kW.

The specifications and configuration of the emitter were the following:

- exposure broadness : 310 mm
  - power: 100 % i.e. 36 kW
  - distance of the smitter to the coating of the plate: 50 mm
  - speed of the place: between 5 and 3 m/minute as indicated in Table 2, 3, 4 and 5.
- Procedure for baking process by static oven-heating, hereinafter referred to as "Static Oven":

[0044] The printing plate precursors TG, TS-1, TS-2 and VPP, as prepared above, were heated in a oven, heated at 235°C during 5 minutes.

Procedure for baking process by dynamic oven-heating, hereinafter referred to as "Dynamic Oven":

[0045] The printing plate precursors TG, TS-1, TS-2 and VPP, as prepared above, were heated with a Haase Oven type OG15 at a temperature setting of 270°C and a speed of 0.7 m/min, corresponding to a dwell-time of the baking step of 150 seconds.

Test for chemical resistance:

9

10

20

[0046] For measuring the chemical resistance, the following corrector solution was selected:

- KP273. Test Corrector Solution for negative and positive thermal plate printing precursor TG, TS-1 and TS-2;
- KN250: Test Corrector Solution for negative violet photopolymer plate printing precursor VPP.

[0047] The chemical resistance was tested by contacting a droplet of the Test Corrector Solution on the baked plates.

After 5 minutes, the droplet was removed from the coating with a cotton pad. The attack on the coating, due to the Test Corrector Solution, was raised by visual inspection as follows:

- Ot no attack,
- 1; changed gloss of the coating's surface,
- 2; small attack of the coating (thickness is decreased).
  - 3; heavy attack of the coating.
- 4: completely dissolved coating.

[0046] The higher the rating, the less is the chemical resistance of the coating. The results for the Test Corrector Solution, hereinafter referred to as "T-C-S", on the coating of each printing plate precursor are summarised in Table 2, 3, 4 and 5. The lables contain also information about the type of the printing plate precursor, the type of the baking process, the speed of the IR-beking process and the dwell-time of the baking process.

Table 2

			580	No S.		
30	Example number	Type printing plate precursor	Type baking step	Speed IR post- baking (m/min)	Dwell time (sec.)	Chemical Resistance (for T-C-S KP273)
33	Comparative example 1	TG	No	<u>.</u>		4
	Comparative example 2	TG	Static Oven	^	300	\$
4D	Comparative example 3	TG	Dynamic Oven	•	150	0
	Example 1	TG	Dynamic IR	5.0	3.7	Q
	Example 2	TG	Dynamic (R	4.0	4.5	0
	Example 3	TG	Dynamic IR	3.5	5.3	9
48	Example 4	TG	Dynamic IR	3.7	5.0	Ď.
	Example 5	TG	Dynamic IR	3,8	4,9	9
	Example 6	TG	Dynamic IR	3.0	6.2	0
		**********		***************************************		

<sup>[0049]</sup> The Examples in Table 2 demonstrate that the baking step, carried out within a dwell-time of less than 7 seconds by a dynamic IR-exposing apparatus, is able to form a TG-printing plate, with an excellent chemical resistance.

55

Table 3:

5	Example number	Type printing plate precursor	Type baking step	Speed IR post- beking (m/min)	Owell time (sec.)	Chemicsl Resistance (for T-C-S KP273)
	Comparative example 4	TS-1	No	-	<del>.</del>	å
10	Comperative example 5	T5-1	Static Oven	٠	390	0
	Comparative example 6	TS-1	Dynamic Oven	-	150	0
	Example 7	TS-1	Dynamic IR	5.0	3.7	Ø
75	Example 6	TS-1	Dynamic IR	4.0	4.8	0
	Example 9	TS-1	Oynamic IR	3.5	5.3	0
	Example 10	TS-1	Dynamic IR	3.7	5.0	©.
20	Example 11	TS-1	Dynamic IR	38	4.9	0
	Example 12	TS-1	Oynamic IA	3.0	8.2	ð

[0050] The Examples in Table 3 demonstrate that the baking step, carried out within a dwell-time of less than 7 seconds by a dynamic IR-exposing apparatus, is able to form a TS-1-printing piece, with an excellent chemical resistance.

Table &

			:8:	No 4:		
30	Example number	Type printing plate precursor	Type baking slep	Speed IR post- baking (m/min)	Owell time (sec.)	Chemical Resistance (for T-C-S KP273)
	Comparative example 7	TS-2	No	•	·	4
33	Comparative example 8	TS-2	Static Oven	-	300	0
	Comparative example 9	TS-2	Dynamic Oven	^	150	0
40	Example 13	TS-2	Dynamic IR	5.0	3.7	0
**	Example 14	TS-2	Dynamic (R	4.0	4.6	9
	Example 15	TS-2	Dynamic IH	3.5	5.3	0
	Example 16	TS-2	Dynamic IR	3.7	5.0	0
45	Example 17	TS-2	Dynamic IR	3.8	4,9	0
	Example 18	TS-2	Dynamic (R	3.0	6.2	0

[0051] The Examples in Table 4 demonstrate that the baking step, carried out within a dwell-time of less than 7 seconds by a dynamic IR-exposing apparatus, is able to form a TS-2-printing plate, with an excellent chemical resistance.

25

Table 5

9	Example number	Type printing plate precursor	Type baking step	Speed iR post- baking (m/min)	Dwell time (sec.)	Chemical Resistance (for T-C-S KN250)
	Comparative example 9	VPP	No	4		å.
10	Comparative example 10	456	Static Oven	~	300	0
	Comparative example 11	VPP	Dynamic Oven	-	150	0
	Example 13	Abb	Dynamic IA	5.0	3.7	Ø
75	Example 14	Abb	Dynamic IR	4.0	4.6	0
	Example 15	Vbb	Oynamic IR	3.5	5.3	0
	Example 16	Abb	Dynamic IA	3.7	5,0	©.
20	Example 17	Abb	Dynamic (B	3.8	4.9	0
	Example 18	Veb	Dynamic IR	5,0	6.2	0

[0052] The Examples in Table 5 demonstrate that the baking step, carried out within a dwell-time of less than 7 seconds by a dynamic IR-exposing apparatus, is able to form a VPP-printing plate with an excellent chemical resistance.

### Claims

25

30

40

- 1. Method for making a lithographic printing plate comprising the steps of
  - providing a lithographic printing plate precursor comprising a photosensitive coating on a substrate having a
    hydrophilic eurlace.
  - image-wise exposing said photosensitive coaling;
  - developing said exposed coating to form an image on said substrate and optionally gumming the plate,
  - baking the image on the plate.

characterised in that said baking step is carried out within a dwell-time of less than 1 minute.

- 2. Method according to claim 1 wherein said baking step is carried out by exposing the printing plate to an infra-red radiation source.
  - Method according to claim 2 wherein said infra-red radiation source is a near infra-red lamp or a middle infra-red lamp.
- Method according to claims 2 or 3 wherein said baking step is carried out by the following steps:
  - radiating by an infra-red source, and
  - transporting said printing plate relatively to said radiation source.
- Method according to any of the preceding claims wherein said photosensitive coating comprises an infra-red absorbing compound.
- Method according to claim 5 wherein said infra-red absorbing compound has an absorption maximum between 750 nm and 3000 nm.
  - Method according to any of the preceding claims wherein said photosensitive coating is an infra-red sensitive coating.

5

10

75

20

25

30

No.

40

48

50

55

8. Method according to any of the preceding claims wherein said printing plate is isolated to prevent heat conduction to the environment. 9. Method according to any of claims 2 to 8 wherein said baking step is carried out by exposing the printing plate to an infra-red radiation source combined with an ultraviolet radiation source. 10. Method according to any of the preceding claims wherein said substrate, having a hydrophilic surface, is an anodised aluminum plate. 11. Apparatus for baking an image obtained after developing an imagewise exposed photosensitive coating on a substrate having a hydrophilic surface, said apparatus comprising an infra-red radiation source, characterised in that said infra-red radiation source is a near infra-red lamp or a middle infra-red lamp. 12. Processor for developing a lithographic printing plate comprising the apparatus according to claim 11. 13. Use of an infra-red radiation source in a method for baking a lithographic printing plate in accordance with ciairn 2, for increasing the speed of the baking process and for obtaining a high chemical resistance and a high printing run length of the plate.



# EUROPEAN SEARCH REPORT

Application Number

EP 03 10 2530

Sategory	Olitation of document with int of relevant possess	sication, where appropriate, es	Fielswant to claim	CLASSIFICATION OF THE APPLICATION UNLCL.T)
A	US 6 040 113 A (VAN 21 March 2000 (2000 * column 7. line 4-1 * column 9, line 11	-03-21) 5 *	1-13	B41C1/10 B41N3/00
Å	EP 1 176 031 A (AGF 30 January 2002 (200 * paragraph [0033] - examples 1-5 *	02-01-30)	1-13	
Å	US 6 362 812 81 (5A) AL) 5 March 2002 (28 * column 8, line 15	VARIAR-HAUCK CELIN ET 902-03-05) - line 17 *	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	***************************************
A	US 6 458 510 B1 (KO) 1 October 2002 (2003 * column 11 *	KKELENBERG DIRK ET AL) 2-10-01)	1-13	***************************************
				TECHNICAL FIELDS SEARCHIO (MCCL.7)
				8410 8418
				W
	·			
	The present search report has b	son drävn up for all alakns.	~	4 0000
	Pure it seems MUNT CH	One of exception of the security 20 November 2003	Spy	ropoulou, E
Ki part Yi part once Ali festi Oli nos	L. NYESORY OF CITED OCCUMENTS industry relevant if taken alone industry relevant if taken alone industry relevant if opmissort with anoth ment of the aeron ordegory relegisted background mediate disclosure mediate document	T theory or principle E service potential E terrice potent do star the Sting dat C document offed i C document offed i E moretaer of the S	s underlying the isomeoni, but public in in the application or other reasons	mistibur ahed on, er

# ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 03 10 2530

This shreek lists the patent family members relating to the patent documents ated in the above mentioned European search report. The members are as contained in the European Patent Office EDP file on

The European Patent Office is in no way liable for these perfouters which are merely given for the purpose of information.

20-11-2003

DE 69808847 D1 05-07-200 DE 69808847 T2 20-09-200 EP 1110720 A2 27-06-200 EP 0864420 A1 16-09-1991 JP 10250255 A 22-09-1991 EP 1176031 A 30-01-2002 EP 1176031 A1 30-01-2001 JP 2002121684 A 26-04-2001 US 2002043168 A1 18-04-2001	DE 6980847 DI 05-07-200 DE 6980847 DI 05-07-200 DE 6980847 T2 20-09-200 EP 1110720 AZ 27-06-200 EP 0864420 AI 16-09-199 JP 10250255 A 22-09-199 DP 10250255 A 22-09-199 DP 2002121684 A 26-04-200 US 2002043168 AI 18-04-200 US 2002043168 AI 18-04-200 DP 2002518715 T 25-06-200 US 6352812 BI 05-03-2002 EP 1011970 AZ 28-06-200 US 6358669 BI 19-03-200 US 6352811 BI 05-03-200 US 635281 BI 05-03-200 US 635281 BI 05-03-200 US 635281 BI 05-03-20		ni port	Publication date	*****	Patent family member(s)	Publication date
JP 2002121684 A 26-04-2003 US 2002043168 A1 18-04-2003 US 6352812 B1 05-03-2002 EP 1011970 AZ 28-06-2004 JP 2002518715 T 25-06-2003 WO 9967097 AZ 29-12-1999 US 6358669 B1 19-03-2003 US 6352811 B1 05-03-2003 US 6534236 B1 18-03-2003 US 6534236 B1 18-03-2003 US 6458510 B1 01-10-2002 EP 1025991 A1 09-08-2003 US 6458510 B1 01-10-2002 EP 1025991 A1 09-08-2003	US 6352812 B1 05-03-2002 EP 1011970 A2 28-06-2001 US 6352812 B1 05-03-2002 EP 1011970 A2 28-06-2001 WC 9967097 AZ 29-12-1991 US 6358669 B1 19-03-2002 US 6362811 B1 05-03-2002 US 6354238 B1 18-03-2002 US 6458510 B1 01-10-2002 EP 1025991 A1 09-08-2001 US 6458510 B1 01-10-2002 EP 1025991 A1 09-08-2001 US 6458510 B1 01-10-2002 EP 1025991 A1 09-08-2001	US 6040113	å	21-03-2000	0E 0E P	69808847 D 69805847 TX 1110720 AX 0864420 AX	1 05-07-200 2 28-09-200 3 27-06-200 1 16-09-199
JP 2002518715 T 25-06-2002 WC 9967097 AZ 29-12-1999 US 6358669 B1 19-03-2002 US 6352611 B1 05-03-2002 US 6534236 B1 18-03-2002 US 6458510 B1 01-10-2002 EP 1025991 A1 09-08-2002 DE 69909733 D1 28-08-2002	JP 2002518715 T 25-06-2002 WC 9967097 AZ 29-12-1999 US 6358669 B1 19-03-2002 US 6352611 B1 05-03-2002 US 6534236 B1 18-03-2002 US 6534236 B1 18-03-2002 US 6458510 B1 01-10-2002 EP 1025991 A1 09-08-2002 DE 69909733 D1 28-08-2002	EP 1176031	A	30-01-2002	38	2002121684 A	26-94-299
DE 69989733 D1 28-08-2003	DE 69989733 D1 28-08-2003	US 6352812	61	05-03-2002	40 20 20 20 20	2002518715 T 9967097 A2 6358669 B1 6352811 B1	25-06-200; 2 29-12-199; 1 19-03-200; 1 05-03-200;
		US 6458510	81	01-10-2002	30	699897 <b>3</b> 3 Di	28-08-200
	more details about this annex : see Official Journal of the European Patent Office, No. 1282						